Claims 1-3, 7-18, 40 and 42-49 remain in this application. Claims 4-6, 19-39 and 41 have previously been canceled without prejudice. Claims 1, 3, 9, 14-15, 40, 42, 44 and 48 currently being amended.

Claim 1 has been amended to specify that the basal medium in step b) is a "first basal medium", and the medium in step c) is a "second basal medium". The terms "first" and "second" have been inserted for the purpose of clarity. Support for this amendment can be found throughout the specification, for example at page 21, bottom paragraph (lines 23-27); page 24, middle paragraph (lines 11-21); page 26, top paragraph (lines 4-11); page 27-28; page 28, paragraph spanning pages 28-29 (page 28, line 24 to page 29, line 2); page 29, bottom paragraph (lines 26-29); Example 5; and Example 6.

Claim 3 has been amended to remove the term "or seedlings", and claim 11 has been amended to specify that the explant is a tissue selected from the group consisting of a seed, petiole, hypocotyl, stem, cotyledon and leaf. Support for the latter amendment can be found throughout the specification, for example page 24, lines 1-2; page 25, middle paragraph (lines 14-16); page 27, middle paragraph (lines 13-15); page 28, middle paragraph (lines 16-18); and page 29, last line to page 30, lines 2.

Claim 9 has been amended to correct a typographical error. Support for this amendment may be found in claim 9 as originally filed. Claim 9 has also been amended to show "-1" in a superscript position.

Claims 14 and 15 have been amended to show "-1" in a superscript position.

Claims 40 and 42 have been amended to depend from claim 1, rather than claim 4. Claim 42 has also been amended to show "-1" in a superscript position.

Claim 44 has been amended to correct a spelling error.

Claim 48 has been amended to correct a clerical error.

For the reasons given below, Applicants submit that the amended claims are in condition for allowance and notification to that effect is earnestly solicited.

Claim Rejections Under 35 U.S.C. § 112

Claim 1 is rejected under 35 U.S.C. 112, second paragraph as being indefinite, as Examiner alleges that it is unclear whether the basal medium from step b) is the same as the basal medium from step c). Examiner further contends that this term is not defined in the specification or claims. Applicant respectfully disagrees with Examiner's objection.

Applicant has amended claim 1 to specify that the basal medium in step b) is a "first basal medium", and the medium in step c) is a "second basal medium". Applicant submits that it would be clear to a person of skill in the art, reading the present specification (e.g., page 21, bottom paragraph (lines 23-27); page 24, middle paragraph (lines 11-21); page 26, top paragraph (lines 4-11); page 27-28; page 28, paragraph spanning pages 28-29 (line 24 to page 29, line 2); page 29, bottom paragraph (lines 26-29); Example 5; and Example 6) that the explant with regenerated tissue can first be cultured on basal medium, then transferred to a fresh, or second basal medium supplemented with an additive of interest (nutrient mineral element) for subculture. Therefore, Applicant submits that the clarifying amendment finds support in the specification.

Furthermore, Applicant submits that the term "basal medium" is defined in the specification on page 21, line 19 to page 22, end of Table 1. Table 1 of the specification provides several examples of basal media that are well known in the art, for example Murashige & Skoog, and Gambourg media. The use of these and similar media (e.g. Linsmaier & Skoog) in the cited prior art (e.g. Stojakowska, Cellarova et al., and Dodds) supports the fact that a person of skill in the art would be familiar with these media. Furthermore, these types of culture media are readily recognized within the prior art as media for plant cell culture, see for example, Potrykus and Shillito (1988), *in* Methods for Plant Molecular Biology, Weissbach and Wessbach (eds), Academic Press, San Diego, pp. 370-373; copy enclosed as Exhibit A).

Therefore, Applicant maintains that a person of skill in the art would understand the meaning of "basal medium", upon reading the present description, and use of their common general knowledge.

Claim 3 is rejected under 35 U.S.C. 112, second paragraph due to an antecedent problem with respect to the seedlings. In response, Applicant has amended claim 3 to remove the expression "or seedlings".

Examiner rejected claim 11 under 35 U.S.C. 112, second paragraph, stating that it is unclear how an explant could be selected from the seed, and that there is no antecedent basis for the seed, petiole, hypocotyls, stem, cotyledon, and leaf. Claim 11 has been amended to specify that the explant is a tissue selected from the group consisting of a seed, petiole, hypocotyl, stem, cotyledon and leaf. Applicant submits that the use of the indefinite article "a" addresses the antecedent problem noted by Examiner. Applicant also maintains that the specification clearly indicates that any such tissues are suitable for use in micropropagation (page 24, lines 1-2; page 25, middle paragraph (lines 14-16); page 27, middle paragraph (lines 13-15); page 28, middle paragraph (lines 16-18); and page 29, last line to page 30, line 2). Furthermore, Applicant points out to Examiner that the use of a seed as an explant is known within the art (see, for example, Smith (1988), *in* Methods for Plant Molecular Biology, Weissbach and Wessbach (eds), Academic Press, San Diego, pp. 343-346, copy enclosed as Exhibit B).

Examiner rejected claims 40 and 42 under 35 U.S.C. 112, second paragraph, stating that they depend from a cancelled claim. In response, Applicant has amended the claims to depend from claim 1, rather than claim 4.

Claim 45 is rejected under 35 U.S.C. 112, second paragraph because the medium contains one or more plant growth regulator, while claim 44 (from which claim 45 depends) specifies a basal medium lacking a plant growth regulator. Applicant respectfully traverses Examiner's objection. Applicant points out to Examiner that claim 45 claims that in the step of culturing (i.e., step a)) of claim 44, the induction medium comprises one, or more than one plant growth regulator. It is submitted that the induction medium of step a) and the basal medium of step b) of claim 44 are two distinct media used in two distinct steps of the claimed method. Therefore, it is maintained that it would be clear to a person of skill in the art that the absence of plant growth regulators in the basal medium does not preclude the presence of plant growth regulators in the induction medium.

Removal of the rejection to claims 1, 3, 11, 40, 42 and 45, and their associated dependant claims under 35 U.S.C. 112, second paragraph is respectfully requested.

Claim Rejections Under 35 U.S.C. § 102/103

Claim 47 is rejected under 35 U.S.C. Section 102(b) as allegedly being anticipated by, or in the alternative under 35 U.S.C. Section 103(a) as obvious over Cellarova et. al. Examiner contends that Cellarova et. al. grows plants in the presence of Ca and Zn, and that the levels of these additives would be elevated when compared to plants grown on a basal medium.

Claim 47 is directed to a phytopharmaceutical plant prepared by the method of claim 1. Claim 47 further defines that the plant of claim 1 comprises an elevated level of the additive of interest "compared to a plant grown in said basal medium in the absence of said additive of interest". Claim 1 specifies that the basal medium is supplemented with from about 50 to about 200 mg/L of an additive of interest selected from the group consisting of a vitamin, boron, chromium, cobalt, copper, iron, lithium, iodine, manganese, molybdenum, nickel, selenium, silicon, tin, vanadium and zinc. Applicant notes that these elements are either micronutrients or a vitamin and are not present in basal media at the amounts indicated. Furthermore, calcium is a macronutrient and not a micronutrient.

Cellarova, (last sentence of page 267) uses a basal medium containing macroelements and microelements according to Linsmaier and Skoog. The composition of the Linsmaier and Skoog (LS) media, along with the composition of other basal media commonly used within the art, was provided in the response to the previous office action. Applicant agrees with Examiner that the LS medium contains calcium and zinc. However, calcium is a macronutrient and is not listed as an additive in claim 1, and zinc is only present as a micronutrient at an amount of 8.6 mg/L.

Cellarova et. al. do not teach or suggest the supplementation of the LS medium with an additive of interest, for example a microelement *in addition to those already present in the medium*. Therefore, Cellarova et. al. do not teach or suggest the supplementation of the LS medium with from about 50 to about 200 mg/L of zinc and growing a plant on this supplemented basal medium.

Applicant submits that Examiner's comparison of a plant grown in a medium containing calcium and zinc to a plant grown on a basal medium lacking these elements is incorrect.

Applicant maintains that, in accordance with claim 47, the amount of additive of interest in a plant grown in basal medium that has been supplemented with about 50 to about 200 mg/L of the

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additive of interest, is to be compared to the amount of additive of interest in a plant grown in *the same basal medium that has not been supplemented*. For example, if the additive of interest were zinc, and the basal medium were LS medium, then the amount of zinc in a plant grown in a media containing LS, supplemented with 50-200 mg/L zinc, would be compared to the amount of zinc in a plant grown in LS. While the plant grown in LS alone would comprise a small amount of zinc, the plant grown in LS plus 50-200 mg/L zinc would comprise a significantly greater amount of Zn. This can be readily seen with reference to Figure 7, and supporting text page 49, middle paragraph (Example 6) of the present application, where plants are grown in basal media ("0" mg/L zinc added to the basal media), and media supplemented by an additional 50 to 200 mg/L zinc. The amount of zinc in plants cultured is shown. A clear 10-fold increase in the levels of zinc can be seen within the plant tissues when the basal media is supplemented with 50 mg/.L zinc, when compared to a plant grown in the basal media alone.

With respect to Examiner's comment that in the absence of any definition of "basal", water could be considered a basal medium, Applicant submits that to one of skill in the art, upon reading the specification would readily understand the term "basal media". As discussed above, Table 1 of the specification provides several examples of basal media that are well known in the art, for example Murashige & Skoog, and Gambourg media. The use of these and similar media (e.g. Linsmaier & Skoog) in the cited prior art (e.g. Stojakowska, Cellarova et al., and Dodds) supports the fact that a person of skill in the art would be familiar with these types of media. These and other culture media are repeatedly identified in the prior art as media suitable for plant cell culture (e.g. Smith (1988), pp 346-7; and Potrykus and Shillito (1988), pp. 367-373, in Methods for Plant Molecular Biology, Weissbach and Wessbach (eds), Academic Press, San Diego; copies enclosed as Exhibits B and A, respectively). Furthermore, Applicant submits that providing that a plant demonstrates an increase in the amount of an additive of interest as defined in claim 47 when compared to a plant grown in the absence of the additive of interest, where the additive of interest is as defined within claim 1 (i.e. the additive of interest is provided at an amount from about 50 to about 200 mg/L, and it is selected from the group consisting of a vitamin, boron, chromium, cobalt, copper, iron, lithium, iodine, manganese, molybdenum, nickel, selenium, silicon, tin, vanadium and zinc) then any media, as would be known to one of skill in the art, may be used as a "basal media".

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There is no teaching, or suggestion in Cellarova et. al. that the basal media is to be supplemented with about 50 to about 200 mg/L, of a compound selected from a vitamin, boron, chromium, cobalt, copper, iron, lithium, iodine, manganese, molybdenum, nickel, selenium, silicon, tin, vanadium and zinc. Therefore, Applicant submits the subject matter of claim 47 is not disclosed or suggested by Cellarova et. al., and removal of the rejection under 35 U.S.C. 102(b), and/or 35 U.S.C. Section 103(a) is requested.

SUMMARY

It is respectfully submitted that the above-identified application is now in a condition for allowance and favorable reconsideration and prompt allowance of these claims are respectfully requested. Should the Examiner believe that anything further is desirable in order to place the application in better condition for allowance, the Examiner is invited to contact Applicants' undersigned attorney at the telephone number listed below.

Respectfully submitted,

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5. The Synthesis of High-Molecular-Weight Proteins in M. the Wheat Germ Translation System G. W.

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[23] Protoplasts: Isolation, Culture, Plant Regeneration By Ingo Potrykus and RAYMOND D. SHILLITO

Protoplasts are isolated, single, and "naked" plant cells. Some have the potential to (1) regenerate a cell wall, (2) dedifferentiate, (3) divide mitotically and proliferate to form unlimited growing cell clones, and (4) differentiate shoot and root meristems (or embryos) which grow out to regenerate complete plants. Protoplasts are thus potentially totipotent individuals at the single cell level. Their freely accessible plasma membrane makes them, in addition, ideal experimental systems for many kinds of genetic manipulation.

The following article is organized in three parts:

1: An idealized example of protoplast isolation will show the steps involved in protoplast isolation and culture.

A general section will discuss the parameters involved in the developmental sequence from protoplast to plant.

3. An experimental section will give a collection of detailed laboratory protocols for protoplast isolation, clonal proliferation, and plant regeneration which are in routine use in independent laboratories, It is recommended that one should gain expertise with one of these easy protocols before approaching a new experimental system. It is not absolutely necessary, but it might be helpful to visit an established protoplast laboratory.

An Idealized Example

An idealized example will be given first to give a general outline of the method of protoplast isolation. For specifics the reader is referred to the detailed protocols.

Plants of a competent genotype of a herbaceous dicot (Petunia hybridg: see also protocol 2 below) are grown in potting compost in clay pots in a controlled environment (12 hr light, 5000 lux, 27/20° day/night temperature, 507/0% relative humidity) and watered with a 0.1% commercial fertilizer solution at 8.00 am and 4.00 p.m.

Leaves expanded to two-thirds their final size are harvested from plants at the 7–9 leaf stage, washed with tap water, and surface sterilized by brief immersion in 70% ethanol followed by 10 min incubation in 0.01% w/v mercuric chloride containing a wetting agent. This is carefully washed off by five rinses each of 5 min with sterile distilled water. The leaves are

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in 10 ml of enzyme solution in 9-cm petri dishes at 28° for ~3 hr. The process of protoplasting is observed from time to time under the inverted microscope. Gentle shaking helps to release protoplasts from the digested volume of enzyme solution and vacuum infiltrated to replace the intercellular air with enzyme solution. Samples of $\sim \! 500$ mg tissue are incubated cut into fine cross sections. These are collected and transferred to a small tissue toward the end of the incubation period.

centrifuge tubes. These are centrifuged (5 min, 60 g) to sediment the The supernatant is carefully pipetted off, the sediment resuspended in the repetitions of this process, the protoplasts (Fig. 1) are taken up in culture medium, a sample taken for counting, and the population density adjusted to $2 \times 10^4/ml$. Aliquots are pipetted into petri dishes to give a thin liquid mixed with an equal volume of osmoticum, and distributed into capped protoplasts without sticking them too tightly to the bottom of the tube. residual 0.5 ml of liquid, and the tube filled with fresh osmoticum. After 2 layer which just covers the bottom. The dishes are sealed with Parafilm The total digest is filtered through a 100-µm stainless-steel sieve, and incubated at a constant temperature of 26° in the dark.

dividing protoplasts in representative fields and calculating the percentage of the total surviving population which has divided at least once. At this reduced osmotic pressure and growth regulator concentration. This is can be estimated 7-10 days after isolation by counting dividing and nonrepeated weekly and after a total of 4 weeks protoplast-derived colonies a size of 1-2 mm in diameter (Fig. 5), the plating efficiency (colonyforming efficiency) is established by calculating how many proliferating Cell wall regeneration and dedifferentiation are visible after one day time the suspension is diluted with 1/3 volume of culture medium with become visible to the naked eye (Figs. 4 and 5). When these have reached (Fig. 2) and the first divisions after three (Fig. 3). The division frequency cell clones have developed per 100 protoplasts originally plated.

solidified medium for further proliferation at low osmotic pressure. After ple shoots (percentage regeneration efficiency) which are rooted as cutings on a medium with a low auxin concentration (Fig. 9). Two weeks later the rooted shoots are carefully washed free of agar, potted into a fine At this stage cell colonies are transferred onto the surface of agara total of 2 months the colonies are large enough (>5 mm in diameter, Fig. 6) to be transferred to regeneration media which promote the develop-After a further month a fraction of the clones will have regenerated multipotting compost, and adjusted slowly to a low relative humidity. The ment of meristems and the outgrowth of shoots (Fig. 7) and roots (Fig. 8) plants can then grow further without any special care (Fig. 10).

[23] PROTOPLAST CULTURE

Parameters Affecting Protoplast Isolation and Proliferation and Plant Regeneration

cussion of these factors will help in understanding the complexity of the phenomenon, and may serve as a guideline to identify mistakes in experiments which fail to reproduce published protocols or where new systems The developmental sequence from protoplast to plant (Figs. 1-10) depends upon numerous parameters and can fail at many points. A disare under study.

Competence

the developmental route from protoplast to plant depends describes the proliferate, and regenerate into a plant is a phenomenon which we can This term is used to describe the ability of a cell to respond in specific The key factor which decides whether or not a protoplast will divide, observe, but about which we have little solid information: competence. capacity of specific plant cells to respond to isolation and in vitro condiways to specific stimuli during development. The competence on which ions with a self-regulating programme of dedifferentiation, proliferation, pattern formation, differentiation, and plant development.

competence when taken into culture cannot be forced to aquire it by in there are numerous cultures and even plants which yield proliferating cally include competence for differentiation and plant regeneration and and physiological basis. We have at present no way to identify competent cells before they respond. Competence can apparently be lost during isolation and culture of protoplasts. We feel that protoplasts which lack This "in vitro competence" probably has a genetic, developmental, vitro manipulation. Competence for sustained division does not automatiprotoplasts from which no differentiation occurs.

may not be a general character of plant cells, but that it is restricted to that all plant cells are totipotent and it may be naive to expect that this is protoplast through to a plant. It is, however, impossible to prove that a The phenomenon of totipotency of somatic plant tissues and isolated cells is well documented. There is, however, no proof for the hypothesis the case. It is easy to prove that plant cells are totipotent by culturing the specific plant cell is not totipotent. Experience suggests that totipotency competent cells.

Plant Species and Genotype

tabacum (Nicotiana, Solanaceae), where it has been relatively easy to There are plant species (and even genera and families), e.g., Nicotiana establish conditions for protoplast proliferation and plant regeneration

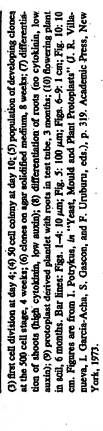
1 W. Halperin, Annu. Rev. Plant Physiol. 26, 395 (1969).

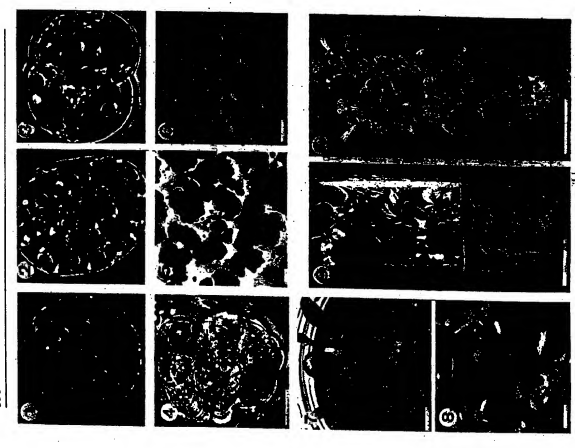
Between these two extremes there are very closely related genotypes of the same species, e.g., Petunia hybrida, where one can routinely regenerate plants from protoplast-derived clones of one genotype and not, so far, from the other. Clear genetic data on this phenomenon are missing and the way out with a nonresponding genotype has been, so far, not so much through variation of culture conditions but rather through screening for responsive genotypes. Genotype screening for in vitro response may pay off in cases where it is not important which genotype is used. Where one must work with a specific genotype, as for example in plant breeding, this option is not available, and one must work on improving other vari-

Source Tissues from Organized Plants and from Cell Culture

Protoplasts are isolated from organized plants and from organized or unorganized cell cultures.

flowers generally give best results. As soon as plants begin to flower, it is difficult, if not impossible, to isolate proliferating protoplasts. The same tions young plants with expanding leaves before the differentiation of fore uses organs before they have fully matured, e.g., expanding leaves at one-half to two-thirds of their final size. Seedling tissues are becoming Developmental state: For plants grown in soil under natural condiholds true for organs where senescence has started. One generally theremore commonly used as a source material. Embryonic or meristematic tissue, or complete embryos, may provide protoplasts with the best poential for proliferation; they are, however, mostly difficult to isolate and





(1) Freshly isolated leaf protoplast; (2) cell wall regeneration and dedifferentiation at day 2; Figs. 1-10. Developmental stages from isolated protoplasts to plant in Petunia hybrida.

concentrations, and long incubation times) may in part interfere with this the isolation procedures required (high osmotic pressure, high enzyme

generally easy to isolate and to handle experimentally and it is usually no Leaf: Leaves are the routine source if no specific reasons make it necessary to use other organs. The reason is that the protoplasts are problem to isolate very large populations of protoplasts at a similar state of differentiation which will proliferate at a high plating efficiency and regenerate plants.

from the epidermis and vascular parenchyma of leaves. It is possible to culture these protoplasts as well. The greater effort required and the lower plating efficiencies obtained are justified, however, only in cases where specific problems require it. This is also true for the culture of the Other differentiated tissues: Parenchymatic protoplasts with small plastids are isolated from stem, petiole, or differentiating root tissues and intensively colored protoplasts isolated from flower petals.

cotyledons, and from seedling roots have been successfully cultured. The protoplasts in cases where attempts to culture protoplasts from leaves Seedlings: These may provide a convenient source of proliferating have failed. Protoplasts from total seedlings, as well as from hypocotyl, disadvantage of limited material may be well compensated for by a proliferation response unobtainable from other organs.

Meristems from the shoot apex or root tips have been used as source in are immature. However, isolation is so tedious and culture so difficult that we cannot recommend this source unless it is absolutely necessary. Embryos: These yield populations of tiny protoplasts as long as they exceptional cases but are not a tissue to be generally recommended be-

easily established and maintained and many successful protoplast labora-Shoot cultures: Axenic shoot cultures proliferating under in vitro concause of reasons similar to those for embryos.

ditions are the source material of choice in species where they can be tures have been established from numerous herbaceous dicots, but not so tories are using them nearly exclusively. There are two types: one is based on normal developed shoots which are rooted and proliferated by rooting shoots under sterile conditions (Nagy and Maliga,2 as described in ing from a wound callus at the base of a shoot grown on a culture meditum with growth regulators (as exemplified by Bindings' work).3 Shoot culfar from any graminaceous species. Protoplasts from shoot cultures are protocol 1); the other is based on multiple adventitious shoots proliferatvery easy to isolate and have a high proliferation and regeneration po-

I. I. Nagy and P. Maliga, Z. Pflanzenphysiol. 78, 453 (1976).
 H. Binding, Physiol. Plant. 35, 225 (1975).

[23] PROTOPLAST CULTURE

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Cell Cultures

entiation is suppressed by a synthetic growth regulator which at the same in an increasing number of plant species, including as interesting groups as the cereals and legumes. The characteristic of these cultures is that small groups of cell have the potential to differentiate to embryos. Differtime enhances proliferation. An embryogenic suspension can be considtinely available in a system such as, for example, carrot. In cereals it has still to be convincingly demonstrated that the embryogenic potential is Organized cell cultures: Embryogenic cultures are becoming available ered an ideal source for totipotent protoplasts and this potential is roupassed through the single cell state.

eral plant species it is possible to establish dedifferentiated cell cultures regulators, with little interference with the morphogenic potential of the Unorganized cell cultures: Cultures capable of regeneration: In sevby enhancing and maintaining the wound response by synthetic growth cells in prolonged culture (e.g., N. tabacum and D. carota). These cultures can be grown on agar-solidified culture media as callus cultures, or can be dissociated into fine suspensions in liquid on shakers. Such culures, from a few suitable genotypes, and with the appropriate treatments, are another convenient source for potentially totipotent protoplasts.

Cell lines: A cell line culture type?) has arisen spontaneously in nearly every plant species kept in culture for long periods of time, including many cereals and grain legumes. Such cell lines are characterized by a nately, a complete lack of morphogenic response. Protoplasts can be easily isolated and, in most cases, cultured. There is, however, very little short cell cycle, proliferation on a single cell basis, friability, and, unfortuchance of plant regeneration from such protoplasts.

Environmental Conditions

yield proliferating protoplast populations. Plants should be watered at plasts can be a key factor. Plants which have been badly treated rarely regular times with the correct amount of water or fertilizer solution. The Care of plants: The care of the plants before the isolation of protosoil type should be appropriate and its structure should provide good acration of the root system. Clay pots are to be preferred to plastic ones. The growth conditions should provide a continuously favorable environment and avoid extremes in any aspect. P. J. Dale, in "Protoplasts 1983: Lecture Proceedings" (L. Potrykus, C. T. Harms, A. Hinnen, R. Hutter, P. J. King, and R. D. Shillito, eds.), p. 31. Birkhauser, Basef, 1983: 5 P. J. King, In "Perspectives in Plant Cell and Tissue Culture" (I. K. Vasil, ed.), p. 25. Academic Press, New York, 1980.

Similar recommendations as for soil-grown plants apply for in vitro

which is grown in incubators without any direct interaction with climatic factors. Experiments which do not work between November and Februa crucial role even with optimized systems. This is not only experienced with greenhouse material, where one can expect it, but also with material Season: It is the experience in many laboratories that season can play ary may work well when repeated between March and June.

before the experiments. The effect of the light quality is less obvious, but Light: This, of course, plays a crucial role. The greatest amount of protoplasts isolated from plants grown under high light intensity are very sensitive to the isolation procedures and, if they survive, proliferate poorly. Experimental plants have therefore to be kept in low light intensity, i.e., 3000 lux or lower, for at least 1 day, and preferably longer, not negligible. It has, however, not been studied carefully enough to allow cool fluorescence tubes should be used in mixtures which cover the visible spectrum more homogeneously than one type of tube. The contribution of the short UV spectrum should be small. If mercury vapor lamps riod probably plays a role too; however, there is no comparative study of are used they should be combined with sodium vapor ones. The photopedata has been accumulated on the effect of light intensity. Generally, clear recommendations. Natural light seems to be superior to artificial this parameter in higher plants.

physiological state of the cell at the time of isolation, but again there are Temperature and relative humidity: These also contribute to the actual no clear data to aid the experimenter.

In general it is advisable to grow the plants under conditions close to

their natural climate.

such a case is often to establish a fresh culture. It should, however, be Care of cell cultures: Cell culture material can be very sensitive to the dilution ratio with fresh culture medium, the material and geometry of the culture vessel, and the aeration and shaking conditions. Isolation of protoplasts is often possible only during a short time period in the exponential growth phase of the culture and quite often cell cultures respond to protoplast isolation totally, or reduces their quality. The only solution in unknown alterations in the culture conditions in a way which prevents made clear that there are many examples of completely unproblematic alterations in the culture conditions, e.g., the time interval of subculture, cell cultures.

Endogenous Factors

within hours, after the physiology of the whole plant in such a way that Response to wounding and pest treatments: Mechanical injury can,

[23] PROTOPLAST CULTURE

protoplast culture is not possible for several days. If, for example, leaves mended that I week of recovery be allowed between experiments. This ments are necessary, there should be a similar time interval between proviso applies for every type of extreme stress situation. If pest treatare taken from the same plant over a long period of time, it is recomtreatment and isolation of protoplasts.

Preculture: Excision of organs and preculture on a culture medium to induce dedifferentiation in situ prior to protoplast isolation has been helpful with many recalcitrant plants, e.g., legumes. On the other hand, attempts at a similar approach with cereal leaves failed completely because the cell walls were modified, in response to even 1 days preculture, such that protoplast isolation was no longer possible.6

Cell cycle phase: The question whether the cell cycle phase plays a role, which phase is optimal for protoplast isolation, and whether actively sively. Fast cycling cells can be sensitive to the isolation procedure but cycling cells are better than resting ones cannot be answered conclucell culture protoplasts are normally isolated from cycling suspension cultures as opposed to resting ones. Protoplasts isolated from fast growing cell suspensions rarely reach the high plating efficiencies of arrested eaf mesophyll protoplasts, but this may be due to other factors than cell cycle phase, such as the state of differentiation.

Sterilization Procedure

The sterilization procedure has to be adapted to the type of organ or the minimum time required for safe sterilization has to be established for pretreatment or to specific sterilizing agents which have to be taken into tissue used in order to ensure that the majority of the cells are not already filled or injured by the sterilizing agent. The minimum concentration and every case. There are also specific sensitivities of tissues to an alcohol account. One of the advantages of sterile in vitro shoot cultures is the fact that there is no danger of oversterilization or incomplete removal of the sterilizing agent.

Mechanical Treatments Prior to Enzyme Application

protect the cells, is not penetrable by the enzymes used for protoplast The outer cell layer of plants has evolved to prevent access to the inner cell layers. The cuttcle, a complex layer of polymers excreted to able are toxic and it is difficult to control their action. The routine proceisolation. It can be degraded by \(\theta\text{-glucuronidase}\), but the enzymes availdure is therefore to mechanically bypass the epidermal cell layer by either pecling off the epidermis (where possible), brushing with an abrasive, or

1. Potrykus, unpublished (1974).

slicing the organ into cross sections. These methods require some care and practice to avoid excessive damage of the inner cell layers.

Plasmolysing Conditions

cells and the intensity of the osmotic pressure used can be critical. This is particularly important for embryonic or meristematic cells which require tinely uses mixtures of mannitol and calcium chloride in the range of 400 to 800 mOsm/kg H₂O. There are, however, cases where these conditions are not successful and where variations in type and concentration of wall from its structural role. Both the agent used for plasmolysing the high osmotic pressures and a long incubation time. Our laboratory rou-Plant cells have an internal pressure (turgor pressure) which is conment would cause the protoplasts to burst. Isolation is therefore carried tained by the rigid cell wall. Digestion of the wall in a hypotonic environout in hypertonic conditions which leads to plasmolysis and frees the cell osmotica have to be screened.

Enzyme Treatment

trolled mixtures of sequences has been possible? but is not required in loses, hemicelluloses, pectins, proteins, and other polymers. Enzymes normally grow on organic matter in forests. The enzymes applied are therefore mixtures of cellulases, hemicellulases, and pectinases. These tion of individual enzymes and application of pure enzymes and conlecular-weight substances is not required in numerous cases where plating for protoplast isolation are commercially available. These are usually obtained from culture filtrates of microorganisms (mostly fungi) which proteases, lipases, phosphatases, and other undesired enzymes. Separamost cases and is therefore not routine. Even purification from low-mo-Plant cell walls are complex chemical and physical structures of cellucommercial enzyme preparations, unfortunately, also contain nucleases, efficiencies higher than 90% can be obtained with unpurified enzymes. However, in more sensitive systems extra purification (by gel filtration) may be essential.

are probably as many variations as laboratories, plants, and types of tinase" at concentrations around 0.5-2% w/v dissolved in an osmoticum (giving 400-800 mOsm/kg H₂O) at pH 5.2-5.6, 24° for a few hours. There The standard treatment is with a mixture of a "cellulase" and a "pecsource material and we can discuss a few general points only.

conditions and growth conditions. Routine established standard protocols Be flexible. The cell walls of plant tissues vary with the environmental

7 S. Ishii and Y. Mogi, in "Protoplasts 1983: Poster Proceedings" (I. Potrykus, C. T. Harms, A. Hinnen, R. Hutter, P. J. King, and R. D. Shillito, eds.), p. 6. Birthauser,

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8

may work perfectly in one laboratory, but fail even if reproduced as precisely as possible in a different laboratory.

The incubation time required for protoplast release depends on the actual state of the source tissue, the enzyme type and mixture, the enzyme concentration, the pH, the osmoticum and osmotic pressure, the incubation temperature, the ratio of tissue: enzyme volume, and the mechanical conditions during incubation.

made concerning the other parameters. The fastest isolation procedure is We have discussed source tissue above. A few comments should be plasts. Efficient enzyme types can be toxic (e.g., pectolyase Y23 or helicase); the optimum mixture has to be established by screening. Embrynot necessarily the best if related to later plating efficiency of the protoonic or meristematic tissues require more hemicellulases and pectinases.

Enzyme concentrations higher than 2% w/v are often deleterious although they reduce the time of incubation. Our laboratory avoids total Plant cells do not normally tolerate pH values below 5.2. Higher pH values and longer incubation times may be necessary with sensitive concentrations above 4% and mixtures of more than 3 commercial enzymes. The pH optimum of nearly all available enzymes is around pH 4.

Mannitol as major osmoticum is sufficient in many cases but there should be always some calcium present (chloride: see protocols) in order to stabilize the plasma membrane. There appears to be no advantage to be gained in using complex osmotica. The osmotic pressure should be as low as possible, but high enough to reduce spontaneous fusion of adjacent cells during isolation.

The incubation temperature should not be raised above 35°. Sensitive tissue be incubated in 10 ml of enzyme solution. It is often essential to systems may benefit from long-term isolations at low temperatures (7-12%). Difficult systems may respond to sequences of high and low temperspoiled many experiments. We recommend that not more than 1 g of incubate under conditions with a large ratio of surface to depth (e.g., 10 atures. The incubation of too much tissue in too little enzyme solution has ml,in a 9-cm petri dish).

the incubation is nearly completed is, however, helpful to maximize protoplast release and shorten the incubation time. Vacuum infiltration of the roller bottles, and with circular shaking of petri dishes on a rocking table or a rotatory shaker. We cannot recommend mechanical shaking during solation of chloroplast-containing cells. Gentle shaking by hand, when Cell culture protoplasts tolerate mechanical shaking during isolation, and in many cases even require this help. There is good experience using enzyme solution into the tissue to replace the intercellular air shortens

treatment the danger lies in the application of a too strong vacuum for too incubation time and improves considerably the homogeneity and quality of protoplast populations isolated from differentiated tissue. With this long a time. Finally it should be mentioned that preplasmolysis for 0.5-1 hr in osmoticum before incubation in enzyme solution can be beneficial.

It requires some experience to recognize, by microscopic examinaplasts can be sensitive to overdigestion as well as to underdigestion. Few tion, the time point when the incubation should be terminated. Protoprotoplasts can recover when the plasma membrane (the plasmelemma) has been disrupted, and "partial protoplasts" having patches of residual cell wall develop far worse than complete protoplasts.

Protoplast Harvest and Purification

plasts. Both washing and purification depend upon differences in the rela-Protoplasts are easily separated from incomplete digested tissues by ive buoyant density of protoplasts and washing solution. Sugar and sugar alcohols (e.g., sucrose and mannitol) have relatively high buoyant densiies. Salt solutions (e.g., CaCl, and MgCl₂) have relatively low buoyant densities at the same osmotic pressure. A practical nonagram giving the relation between osmalility, molality, and buoyant density has been pubished.8 Chloroplast-containing leaf protoplasts have high buoyant densities, cytoplasmic protoplasts have medium, and vacuolated parenchy-Washing and collection by sedimentation require the combination of prolively low buoyant densities. For flotation of protoplasts the opposite applies. Care must be taken to maintain a similar osmotic pressure sieving through meshes with pore sizes adjusted to the size of the protomatic protoplasts with small plastids have low buoyant densities. oplasts with a higher buyoant density with washing solutions with relathroughout the procedures and a convenient osmometer is of great help. Protoplasts should be washed at least three times by repeated sedimentaion or flotation and resuspension. A flotation step may be required for purification in cases where the protoplasts are contaminated with debris which does not separate out during the sieving step. If this is not possible then swirling of protoplasts sedimenting in a low density osmoticum in petri dishes helps to collect the perfectly spherical protoplasts in the center. Step gradients have been described which allow not only purificaferent buoyant densities, which may correlate with their capacity for ion but also fractionation of protoplast populations into fractions of difproliferation. Gradients of osmotically inactive substances such as Per-

C. T. Harms and I. Potrykus, Theor. Appl. Genet. 53, 57 (1978).

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coll9 in combination with osmotica of low buoyant density may yield clean populations where simple osmotica alone fail. Finally, automated cell sorting is also possible with protoplasts, although this is not yet sufficiently developed to be in routine use.

Careful separation of protoplasts from the enzymes used for isolation is essential, as resynthesis of a cell wall, which is inhibited by residual enzymes, is normally an absolute prerequisite for proliferation.

Protoplast Culture

here are numerous further factors which could interfere with or promote Among these are the population density, the composition of the culture Assuming that there was no important mistake in the procedure so far, the capacity of the isolated protoplast to dedifferentiate and to proliferate. medium, particularly growth regulators, the osmotic pressure, physical culture conditions, and undefined factors in the culture medium.

divide at all outside a specific population density range. It is possible to vidual single protoplasts in micro drops 10 which suggests that the ratio of ing efficiency with regard to population density, and protoplasts do not protoplast population densities is between 2×10^4 and 2×10^5 ml, ingrow optimized systems down to very low population densities and indiprotoplasts to volume of culture medium is important. The usual range of Population density: This is a clear case and can be tested easily. Protoplast populations, with exceptions, express a clear optimum of platcreasing with decreasing protoplast size and vice versa.

important role. This typically contains ions of inorganic salts, vitamins, a carbon and energy source (sucrose or glucose), growth regulators, and an osmostabilizer (see the table). No protoplast culture medium has been completely optimized by checking the effect of variations in every single Culture medium: The composition of the culture medium plays an component against every other component; this is also not required, as mized media. There are efficient screening techniques available such as the "Microdrop-Array Technique" 11.12 which can be used to study the effect of systematic variations of culture medium factors. If changes in population density and the established culture media fail to induce sustained divisions the first parameters to vary are the growth regulators. plating efficiencies of higher than 90% are obtained using partially opti-With the culture medium giving best survival and a fixed population den-

W. Wernicke, H. Loerz, and B. Thomas, Plant Sci. Lett. 15, 239 (1979).

¹⁰ K. N. Kao, Mol. Gen. Gener. 150, 225 (1977).

¹¹ I. Potrykus, C. T. Harms, and H. Lourz, Plant Sci. Lett. 14, 231 (1979a).
¹² C. T. Harms, H. Loerz, and I. Potrykus, Plant Sci. Lett, 14, 237 (1979).

sity of 2-5 imes 104/ml, a combination of a dilution series of an auxin versus followed by similar combinations with other types of auxins and cytokinins used in the first screen. Combinations of different auxins (e.g., 2,4-D and NAA) may help. It may be necessary to reduce the hormone a cytokinin both in the range between 0.05 and 5 mg/liter should be tested concentrations by up to 10-fold after a few days of culture.

however, important to make the layer of protoplast suspension as thin as possible (e.g., 0.7 ml in a 3.5-cm petri dish; 2 ml in a 6-cm petri dish). The Most frequently "liquid thin layer" cultures in petri dishes will do. It is, depth of the liquid layer alone can inhibit proliferation. Some protoplasts develop only if plated into a gel, others do not develop under these condibut agarose gives better plating efficiencies and enables development of The physical culture conditions are a further important parameter. tions. Agar as gelling agent may suffice with insensitive protoplast types, ments of protoplast containing gels are floated in larger volumes of liquid sensitive protoplasts. The "bead-type" culture technique where segculture medium and acrated on a rotary shaker has considerably improved the plating efficiency in many systems and has enabled sustained development in systems where protoplasts did not previously divide. 13

Microdrops of 20-40 µl are helpful if the total number of protoplasts is limited. 10. A culture medium with a high buoyant density which maintains the protoplasts floating during the critical initial culture stages, or a floating membrane or filter support onto which the protoplasts are plated are further possible variations.

dium can be sufficient to inhibit development. It is important to check this early in the experiments by culture in a series of different osmotic values tions. Unfortunately, not only the quantity of the osmostabilizer but also its quality are important and there are even interacting effects of the type Osmotic pressure: An incorrect osmotic pressure of the culture medown to $\sim \! \! 300 \, \mathrm{mOsm/kg \, H_2O}$, and to recheck its effect with altered condiculture medium should be gradually reduced to lower and lower levels as of osmoticum used during isolation and the type of culture medium used for subsequent culture. The relatively liigh osmotic pressure of the initial the culture develops.

Environmental conditions: Of the environmental factors temperature ommended as a baseline. There are, however, protoplasts which will not and light are most important. A continuous temperature of 24-26° is recdivide unless cultured at 28° or higher, some of which will not develop at this temperature, and others which require a lower temperature. An initial phase of 12-24 hr at 12° has been found to promote subsequent develop-

ment. Fluorescence tubes are a suitable light source. The light quality ment at 24° in some cases. Light is normally not required for induction of divisions, although it may enhance further development. The light intenplays a role as has been found in experiments with monochromatic light. sity should not exceed 1000 lux, as high intensities inhibit early develop-There are, however, no clear and conclusive data available.

The choice of container in which to culture the protoplasts can have profound effects. Plasticware is now used in most laboratories and a range of petri dishes from different suppliers should be used. We have ourselves found that some brands (e.g., Falcon) can be deleterious under our conditions to even robust protoplasts such as those from N. tabacum.

Undefined factors: There is a further group of factors to be tested if all the variations suggested above do not succeed: conditioned medium. nurse cultures and feeder layers, plant extracts, and X-plates.

sions and added to culture media in different proportions. The time point for the harvest is critical and recommended to be during the early log Conditioned medium is harvested from actively dividing cell suspenphase of growth.

cultures into the surrounding culture medium by coculturing protoplasts Nurse cultures make use of diffusible substances released from cell and cell cultures. The growth of the cell cultures in this combination may inhibited prior to plating by X-irradiation. 14 Another approach to feeding be inhibited by the high osmotic pressure required for the protoplasts. pendent layer (usually below the protoplast layer); often the cells of the feeder layer are protoplasts with high division potential which have been is the use of competent auxotrophic protoplasts which are mixed and Reeder layers consist of protoplasts or cells plated separately in an indeplated with the nondividing protoplasts and later selected against by nonpermissive conditions. 15

X-plates, where a petri dish is separated into quadrants by permeable bars and protoplasts are plated atternatively with culture medium with a lower osmotic potential16 or containing activated charcoal,17 has been reported to improve or enable protoplast division. The authors have no personal experience with this culture technique.

¹³ R. D. Shillito, J. Paszkowski, and I. Potrykus, Plant Cell Rep. 2, 44 (1983).

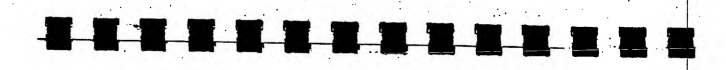
¹ D. Raveh, B. Huberman, and E. Galun, In Vitro 9, 216 (1973).

¹⁵ T. Hein and O. Schieder, in "Protoplasts 1983: Poster Proceedings" (J. Potrykus, C. T. Harms, A. Hinnen, R. Hutter, P. J. King, and R. D. Shilito, eds.), p. 106. Birthauser, Basel, 1983.

stein, B. Regenbach, R. L. Phillips, and C. E. Green, eds.), p. 185. Univ. of Minnesota "J. F. Shepard, in "Genetic Improvement of Crops: Emergent Techniques" (I. Ruben-Press, Minneapolis, 1989.

⁷ I. Carlberg, K. Gilmeilus, and T. Erikason, is "Protoplasts 1983: Poster Proceedings" (L. Potrykus, C. T. Harms, A. Hinnen, R. Hutter, P. J. King, and R. D. Shillito, eds.), p. 258.

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Vitamins and other organics used in culture media (mg/liter concentration)

သ	E bas W	BSB	යා	ေ	K0,-	•••1/H	EZI.	r2		
06	100	1000	100.00	100.00	00,001	100.00	100.00	00.001	100.00	jotizonI-m
			10.0	• •	10.0	10.0			80.0	*saines!\ Biotin
				08.0	00.1	00.I	01.0		05.0	Pyridoxine-HCI
0.1	1.0	1.0	00.1		00.01	10.00	00.1	40.0	05.0	DH-annaydT
2.8	1.0	0.1	10.00	01.0	1.00	1.00				Nicotinamide
		• • •	1.00	0.50			01.0	. •	00.2	Nicotinic acld
· 0°9	2.0	1.0	VFU	مصم	. 05.0	01.0			. 02.0	Folic acid
			04.0		1,00	1.00	•			D-Cs-pantothenate
			00.1 20.0		20.0	20.0				p-Aminobenzoic acid
			00.1		00.1	00.1		•		Choline chloride
•			02.0		02.0	02.0				Ribodavin
•	•	•	2.00		2.00	2,00		٠.		Ascorbic acid A nimetiV
•		•	10.0			10.0 ·		•	•	Vitamin D.
	•		10.0		10.0	10.0		•		Vitamin B ₁₂
•			20.0	•	20.0	20.0			2.00	Olycine
2.0	0°E .	•				01.0				Coconnt water (%4/4)

CULTURE

2

^{*} Where the inorganic component is common, i.e., K3 and CPW media and HAFEO, these have been given together:

• J. P. Mitsch and C. Mitsch, Jetence 163, 85 (1969).

^{*} B. M. Linemaier and P. Skoog, Physiol. Plant. 18, 100 (1965).

I. I. Nagy and P. Maliga, Z. Phoneomphysiol, 73, 653 (1976).
 K. M. Kao and M. R. Michaylai, Planta 126, 105 (1975).

T. Murathigo and F. Skoog, Physiol. Plant. 15, 473 (1962).

S.-H. Xim, M. R. Davoy, and B. C. Cocking, Z. Plantemphylol. 194, 289 (1981).

L. Gemborg, R. A. Miller, and R. Obysma, Exp. Cell Res. 59, 151 (1968).
 H. M. Wood and A. C. Bramp, Proc. Natl. Accel. Sel. U.S.A. 47, 1907 (1961).
 J. Fottylens, C. T. Haims, and H. Loerz, Theor. Appl. Genet. 54, 209 (1979).

^{*} Macroelements are nausity made up as a 10% concentrated stock solution, and microelements are a 1000% concentrated stock solution.

* Citric, furnante, and maife acid (each #0 mayliter final coine.) and sodium pyruvate (20 mayliter) are prepared as a 100% concentrated stock solution, adjusted to pH 6.5 with NH,OH, and added to these media.

Adenine (0.1 mg/liter), and guanine, thyimine, uracil, hypoxanthine, and cytosine (0.00 mg/liter) are prepared as a 1000× concentrated stock solution, adjusted to pH 6.5 as above and added to these media.

isoleucine, leacine, lysine, methionine, phenylalanine, proline, serine, threonine, tryptophen, tyrosine, and valine (0.1 mg/liter). * The following amino scids are added to this medium using a 10× stock solution (pH &5 with NH,OH) to yield the given final concentrations: glutamine (5.6 mg/liter), atamic scid (0.6 mg/liter), cysteins (0.2 mg/liter) asparagine, aspartic acid, cystine, histidine,

Vitamin stock solution is normally prepared 100× concentrated.

Finally, extracts of various kinds may help if everything else has and xylem exudate have improved response in early phases of studies and failed. Coconut water, leaf extracts, extracts from meristematic cultures, could later be replaced by completely defined culture media.

Plant Regeneration

Plant regeneration from cell cultures derived from complex explants may present a completely different problem to plant regeneration from protoplast-derived cultures. Explant-derived cell cultures may, and often do, contain adventitious meristems which proliferate as meristems. Regeneration is then not more than a release of the morphogenic potential of and are able to maintain it over many subcultures, even if it is possible to the preexisting meristems by removal of some sort of repression. Protoplast-derived cell clones have to build new meristems in a mass of dedif. ferentiated cells and there is no guarantee that cells have this potential regenerate plants from complex explants. Fortunately, many genotypes of herbaccous plant species have this inherent capacity (competence for plant regeneration from isolated single cells). In those cases where the cells are competent, plant regeneration is easy, and with the best model plants also very efficient. A relative increase in cytokinins and decrease in auxins in the culture medium, and the presence of light, are the basic duction, maturation, and shoot differentiation to proceed. Low levels of conditions which allow the self-regulating process of shoot meristem inauxins in the absence of cytokinins in the culture medium favor the development of roots. In a few cases conditions have been discovered which allow the development of somatic embryos and their later maturation and germination. ¹⁸ In cases where the inherent competence is missing we are,

Concluding Remarks

Our interpretation of the biological phenomenon of sustained development and plant regeneration from isolated protoplasts is that competent cells have a self-regulatory program to undergo dedifferentiation, susa "wound response" initiated via isolation and amplified by hormone tained mitotic divisions, and differentiation. This program is triggered by treatments. The experimenter helps by providing nutritional and environmental conditions which allow this process to proceed. This works with competent cells and does not work with noncompetent cells. Formation of shoot and root menistems from protoplast-derived cultures depends again on this biological phenomenon of competence, which we can observe, but not define.

³ L. Li and H. W. Koblenbach, Plant Cell Rep. 1, 209 (1982).

[23] PROTOPLAST CULTURE

Protocols for the Preparation and Culture of Protoplasts and Regeneration of Plants We describe four protocols which are in everyday use in our laboratory and one which has been described in the literature.

Centrifugations are carried out at 60 g (100 g in protocol 3) except A number of things are common to these protocols.

where otherwise stated.

Washing solutions (osmoticum) in protocols 1, 2, 4, and 5 are buffered with 0.5% w/v 2-(N-morpholino)ethanesuffonic acid (MES) and adjusted to pH with KOH except where otherwise stated.

ing a drop of a 1:10 dilution of the suspension in calcium chloride in a hemocytometer, counting, and estimating the density in the original sus-Counting of protoplasts: counting of protoplasts is carried out by plac-

Abbreviations

MES: 2-(N-morpholino)ethanesulfonic acid

NAA: naphthaleneacetic acid

2,4-D: 2,4-dichlorophenoxyacetic acid

p-CPA: p-chlorophenoxyacetic acid

BAP: 6-benzylaminopurine

Materials

Sources for the plant material are given with each individual pro-

Table top centrifuge: Universal 2 (Hettich Centrifugen, 72, Tutlingen, West Germany).

Osmometer: Roebling Micro-Osmometer (Infochroma AG., Baarerstrasse 57, CH-6300, Zug, Switzerland)

Rocking table: Heidolph Reax 3 rocking table (Salvis AG., CH-6015, Reussbuehl. Switzerland).

The 10-cm-diameter containers used for the "bead-type" culture are obtained from Semadani AG (Ostermundigen, CH-3072, Switz-Stainless steel sieves: Saulas and Co. (St. Louis, Paris 10, France). erland.)

Petri dishes: these are obtained from a range of suppliers but we have found that those from some suppliers can be toxic to some protoplast types.

SeaPlaque agarose: Marine Colloids, FMC Corp. (5 Maple Street, Rockland, Maine 04841).

Cleaned agar: this is prepared by washing with water, acetone, and

ethanol in succession. 13,19

(ween 80: ICI (Runcorn, England) or Merck-Schuchardt (Hohen-

brunn, Munchen, West Germany)

Percoll: Pharmacia (Uppsala, Sweden).

Cellulase "Onozuka" R10 and Macerozyme R10: Yakult Pharmaceu-Greenzit, Ciba-Geigy AG (Basel, Switzerland).

tical Ind. Co. Ltd. (Shingikan Cho, Nishinomiya, Japan).

Meicelase: Meiji Seika Kaisha Ltd. (Tokyo, Japan).

Driselase: Chemische Fabrik Schweizerhalle (CH-4013, Basel, Switzerland)

Hemicellulase: Sigma Chemical Co. (P.O. Box 4508, St. Louis, Mo. 63178).

Rhozyme HP150: Roehm and Haas Co. (Philadelphia, Penn.).

Pectinol: Roehm GmbH Chemische Fabrik (Darmstadt, West Germany).

All other organic and inorganic substances were of the highest purity available from the usual commercial sources.

1. Preparation and Culture of Protoplasts from a Sterile Shoot Culture, and Regeneration of Plants

used genotype of N. tabacum cv. Petit Havana, SR1.20 This material is The example given is for protoplasts from shoot cultures of the widely grown as sterile axenic shoot cultures.

The protocol for protoplast isolation is modified from that of Nagy and house grown material. In the latter case the leaves must first be surface sterilized (see protocol 2) and then either sliced or the lower epidermis removed by peeling. The culture method uses complex media based on system described by Shillito et al. 13 to obtain high division frequencies Maliga2 and works equally well with tobacco leaf material from greenthat of Kao and Michayluk21 (see the table) and the agarose "bead-type" and rates of conversion of protoplasts to colonies.

Colonies are transferred to agar-solidified medium for 1 subculture and then placed on regeneration medium to promote the formation of shoots. Regenerated shoots are cultured on the original shoot culture medium.

Source of Material. Shoot cultures are established from seed which is sterilized using mercury chloride (see protocol 2) or sodium hypochlorite (5 min, 1.4% w/v containing 0.05% w/v Tween 80) and plants arising are

¹⁰ A. C. Brann and H. N. Wood, Proc. Natl. Acad. Sci. U.S.A. 48, 1776 (1962).

" P. Maliga, A. Sz. Breznovits, and L. Marton, Nature (London), New Biol. 244, 29 (1980).

²¹ K. N. Kao and M. R. Michayluk, Planta 126, 105 (1975).

serially subcultured every 6 weeks as cuttings on T medium2 (see the table) solidified with 0.8% w/v cleaned agar at 26° in 16 hr/day light (1000–

2000 lux) in a growth chamber.

floated on enzyme solution (1.25% w/v Cellulase "Onozuka" R10, 0.4% w/v Macerozyme R10 in K3A medium with 0.4 M sucrose) in petri dishes Preparation of Protoplasts. Just fully expanded leaves of 6-week-old shoot cultures are removed under sterile conditions and wet thoroughly with enzyme solution. The leaves are then cut into 1- to 2-cm squares and (~1 g leaves in 12 ml enzyme solution in a 9-cm diameter petri dish). These are sealed and incubated overnight at 26° in the dark.

The digest is gently agitated and then left for a further half hour to steel mesh sieve and washed through with one-half volume of 0.6 M sucrose (MES, pH 5.6), distributed into capped centrifuge tubes, and complete digestion. The solution is filtered through a 100-µm stainlesscentrifuged for 10 min.

dium is then removed from under the protoplasts. A simple method of doing this uses a sterilized canula (A. R. Howell Ltd., Kilburn High Rd., This must be done slowly so as to avoid disturbing the layer of protoplasts excessively. Alternatively, the protoplasts can be collected using a pi-The protoplasts collect at the upper surface of the medium. The me-London NW6, England) attached to a 60-ml disposable plastic syringe. pette (with a medium orifice).

The protoplasts are resuspended in K3A medium (see the table) containing 0.4 M sucrose. Washing of the protoplasts is carried out by repeated $(3\times)$ flotation and replacing of the medium in this way. A sample is taken for counting before the last centrifugation, and the protoplasts resuspended the last time in H medium (see the table), at a concentration of 10° to 2 × 10°/ml.

Atternatively the protoplasts can be resuspended in K3A or other media or osmoticum and manipulations such as transformation or fusion carried out (e.g., as described by Paszkowski and Saul²³).

Culture. The protoplast suspension in H medium is pipetted into petri ume of liquified K3A medium (0.3 M sucrose, Nagy and Maliga,2 see the dishes (1.5 ml/6-cm-diameter petri dish, respectively) and an equal vol-The protoplasts are distributed evenly by gentle swirling, and the medium table) containing 1.2% w/v ScaPlaque agarose, cooled to 40°, is added. allowed to gel.

the dark at 26° overnight, followed by transfer to the light (500 lux) at the Petri dishes are sealed with Parafilm or an equivalent and incubated in same temperature for a further 6 days.

2 J. P. Nitsch and C. Nitsch, Science 163, 85 (1969).

D J. Paszkowski and M. W. Saul, this volume [28].

;

Cleaned agar: this is prepared by washing with water, acctone, and ethanol in succession. i3.19

Ween 80: ICI (Runcorn, England) or Merck-Schuchardt (Hohenbrunn, Munchen, West Germany).

Percoll: Pharmacia (Uppsala, Sweden).

Greenzit, Ciba-Geigy AG (Basel, Switzerland).

Cellulase "Onozuka" R10 and Macerozyme R10: Yakult Pharmaceu-

tical Ind. Co. Ltd. (Shingikan Cho, Nishinomiya, Japan).

Meicelase: Meiji Seika Kaisha Ltd. (Tokyo, Japan).

Driselase: Chemische Fabrik Schweizerhalle (CH-4013, Basel, Switzerland)

Hemicellulase: Sigma Chemical Co. (P.O. Box 4508, St. Louis, Mo.

Pectinol: Roehm GmbH Chemische Fabrik (Darmstadt, West Ger-Rhozyme HP150: Roehm and Haas Co. (Philadelphia, Penn.).

All other organic and inorganic substances were of the highest purity available from the usual commercial sources. many).

1.. Preparation and Culture of Protoplasts from a Sterile Shoot Culture, and Regeneration of Planis :

used genotype of N. tabacum cv. Petit Havana, SR1.20 This material is The example given is for protoplasts from shoot cultures of the widely grown as sterile axenic shoot cultures.

house grown material. In the latter case the leaves must first be surface sterilized (see protocol 2) and then either sliced or the lower epidermis removed by peeling. The culture method uses complex media based on The protocol for protoplast isolation is modified from that of Nagy and that of Kao and Michayluku (see the table) and the agarose "bead-type" system described by Shillito et al. 13 to obtain high division frequencies Maliga2 and works equally well with tobacco leaf material from greenand rates of conversion of protoplasts to colonies.

Colonies are transferred to agar-solidified medium for 1 subculture and then placed on regeneration medium to promote the formation of shoots. Regenerated shoots are cultured on the original shoot culture medium.

sterilized using mercury chloride (see protocol 2) or sodium hypochlorite (5 min, 1.4% w/v. containing 0.05% w/v Tween 80) and plants arising are Source of Material. Shoot cultures are established from seed which is

7 A. C. Braun and H. N. Wood, Proc. Natl. Acad. Sci. U.S.A. 48, 1776 (1962).

D. Maliga, A. Sz. Breznovits, and L. Marton, Nature (London), New Biol. 244, 29 (1980). ²¹ K. N. Kao and M. R. Michayluk, Planta 126, 105 (1975).

· [23] PROTOPLAST CULTURE

serially subcultured every 6 weeks as cuttings on T medium2 (see the table) solidified with 0.8% w/v cleaned agar at 26° in 16 hr/day light (1000–

2000 lux) in a growth chamber.

Preparation of Protoplasts. Just fully expanded leaves of 6-week-old with enzyme solution. The leaves are then cut into 1- to 2-cm squares and floated on enzyme solution (1.25% w/v Cellulase "Onozuka" R10, 0.4% w/v Macerozyme R10 in K3A medium with 0.4 M sucrose) in petri dishes shoot cultures are removed under sterile conditions and wet thoroughly (~1 g leaves in 12 ml enzyme solution in a 9-cm diameter petri dish).

The digest is genily agitated and then left for a further half hour to These are sealed and incubated overnight at 26° in the dark.

steel mesh sieve and washed through with one-half volume of 0.6 M sucrose (MES, pH 5.6), distributed into capped centrifuge tubes, and complete digestion: The solution is filtered through a 100-µm stainlesscentrifuged for 10 min.

This must be done slowly so as to avoid disturbing the layer of protoplasts excessively. Alternatively, the protoplasts can be collected using a pidium is then removed from under the protoplasts. A simple method of doing this uses a sterilized canula (A. R. Howell Ltd., Kilburn High Rd., The protoplasts collect at the upper surface of the medium. The me-London NW6, England) attached to a 60-ml disposable plastic syringe. pette (with a medium orifice).

peated $(3\times)$ flotation and replacing of the medium in this way. A sample is taken for counting before the last centrifugation, and the protoplasts resuspended the last time in H medium (see the table), at a concentration of The protoplasts are resuspended in K3A medium (see the table) containing 0.4 M sucrose. Washing of the protoplasts is carried out by re- 10^5 to 2 × 10^3 /ml.

media or osmoticum and manipulations such as transformation or fusion Atternatively the protoplasts can be resuspended in K3A or other carried out (e.g., as described by Paszkowski and Saul²³).

ume of liquified K3A medium (0.3M) sucrose, Nagy and Maliga, 2 see the Culture. The protoplast suspension in H medium is pipetted into petri dishes (1.5 ml/6-cm-diameter petri dish, respectively) and an equal vol-The protoplasts are distributed evenly by gentle swirling, and the medium table) containing 1.2% w/v ScaPlaque agarose, cooled to 40°, is added. allowed to gel.

Petri dishes are sealed with Parafilm or an equivalent and incubated in the dark at 26° overnight, followed by transfer to the light (500 hux) at the same temperature for a further 6 days.

2 J. P. Niuch and C. Nitsch, Science 163, 85 (1969).

n J. Paszkowski and M. W. Saul, this volume [28].

The agarose gel containing the protoplast-derived cells is cut into quadrants and transferred to 30 ml of liquid medium (1:1 mixture of K3A and H460) in 10-cm-diameter containers. These are placed on a gyratory

After 1 week one-half of the medium is replaced with a 1:1 mixture of shaker (80 rpm, 1.25 cm throw) in the dark at 26°.13

be split into 2 containers after 3 weeks and again after 5 weeks where no replaced weekly with a 1:1 mixture of K3E and J media. Cultures should K3C and I media (see the table), and thereafter one-half the medium is selection regime is being employed, as otherwise there will be too many colonies per container to allow full development.

diluting the culture weekly with the media described or embedding into Should it be necessary to culture the protoplasts in liquid medium in K3A medium or a 1:1 mixture of this medium and H460 as used above, agarose or agar after I week (see Paszkowski and Saul²³) or at a later time. (e.g., after fusion treatments), they can be cultured at 5 imes 104-105 per ml

Regeneration of Plants. Colonies of 1-2 mm diameter are taken with forceps and placed on Linsmaier and Skoog24 medium solidified with 0.8% w/v cleaned agar (L.S., see the table) and incubated in the dark at 26°.

After 3-5 weeks, depending on the size of the original colony, these should reach 1 cm in diameter. Each colony is then split into 4 parts and $\mathbf{2}$ placed on fresh LS medium as above, and 2 on LS medium with $0.3~\mathrm{mg/}$ liter BAP as the only phytohormone (regeneration medium). These latter dishes are incubated in the dark for 1 week and thereafter in the light (3000-5000 lux).

Shoots arising from the callus on regeneration medium are cut off when 1-2 cm long and placed on LS medium as above, but without hormones. When the shoots reach 3-5 cm in length they are transferred to T medium and treated as shoot cultures (see above).

system: the agar is gently washed away and the plantlets potted up. They They can be transferred to soil once they have an established root require a humid atmosphere for the first week and can then be hardened off and grown under normal greenhouse conditions.

2. Preparation and Culture of Leaf Mesophyll Protoplasts from Greenhouse Grown Plants (Petunia hybrida)

The method described is based on that of Durand et al.25 as described by Shillito et al. 13 and further modified by S. Kruger-Lebus²⁶ in our laboratory for use with the "cyanidin type" and other petunias. A high

[23] PROTOPLAST CULTURE

division frequency and rate of conversion to colonies of protoplasts from

a number of Petunia species is achieved by this method

cles are clonally propagated via cuttings, grown in clay pots in a controlled environment [12 hr light (5000 lux), 27/20° day/night, 50/70% related Source of Material. Plants of Petunia hybrida and other Petunia spoive humidity] and watered morning and evening with commercial fertilizer (0.1% v/v Greenzit).

Preparation of Protoplasts. Young leaves at approximately two-thirds of their final size are washed with tap water and sterilized by immersion for 10 sec in 70% v/v ethanol followed by 10 min in 0.01% w/v HgCl. solution containing 0.05% w/v Tween 80, and then washed carefully with 5 changes of sterile distilled water (each change 5 min).

tol, 0.04 M CaCl₂, MES, pH 5.8) and arranged in a stack of 6 on the lid of a 9-cm petri dish ready for cutting. They are cut carefully into clean sections 0.5 mm wide, transferred into a small screw top flask containing 10 ml of enzyme solution (2% w/v Cellulase "Onozuka" R10, 1% w/v Leaf halves without midribs are wet with osmoticum P1 $(0.3 M \, \mathrm{manni})$ hemicellulase and 1% w/v pectinol in osmoticum P1), and vacuum infiltrated until the leaf tissue is translucent.

The leaf slices are placed in fresh enzyme solution in a petri dish (0.5 at 28° for ~3 hr. The incubation mixture is checked periodically under the inverted microscope for the release of protoplasts. The time required may g/10 ml in a 9-cm petri dish), which is sealed with Parafilm and incubated vary, especially with greenhouse grown material.

tubes. Osmoticum P2 (0.375 M mannitol, 0.05 M CaCl., MES, pH 5.8) is The digest is gently agitated; filtered through a 100- \mm mesh stainlesssteel sieve, and transferred in 5-ml aliquots into 10-15 ml centrifuge added (5 ml) to each tube and, after gentle mixing, these are centrifuged or 5 min to sediment the protoplasts.

P2. Washing by sedimentation is repeated 2 times. If necessary, the suspension is overlaid on 0.6 M sucrose to remove debris and the protoplasts The supernatant is carefully pipetted off, and the sediment is gently shaken to free the protoplasts before resuspension in 10 ml of osmoticum collecting at the interface recovered and resuspended in osmoticum P2. A sample is taken and diluted for counting and the protoplasts sedimented once more and resuspended in medium (K0, see the table) at 10^5 to $2 \times$

Culture. The suspension is pipetted into 9-cm-diameter petri dishes (3 ml per dish) and 3 ml of liquefied K0 medium containing 1.2% w/v Sca-Plaque agarose at 40° added. The protoplasts are distributed evenly by gentle swirling and the medium allowed to set.

The cultures are incubated for 6 days in the dark at 26°. Half the

^{21.} Durand, I. Potrykus, and G. Donn, Z. Pflanzenphysiol. 69, 26 (1973). M. E. M. Linsmaier and F. Skoog, Physiol. Plant. 18, 100 (1965).

⁷⁰ D. Hess, Planta 59, 567, (1963)

These are incubated at 26° on a gyratory shaker (80 rpm, 0.6 cm throw) in the light (500-1000 lux). The liquid medium is replaced weekly, each time diameter containers each containing 40 ml of liquid K0 medium containing reducing the glucose concentration in the original medium by one-quarter protoplast-containing againse gel is transferred to each of two 10-cma one-tenth concentration of the hormones and with 2% v/v coconut milk. so as to reduce the osmotic pressure.

After 6-8 weeks the colonies are 2-3 mm in size and can be cultured further.

tion medium (NT; see the table) with 0.8% w/v ScaPlaque agarose, 2% w/v sucrose, and I mg/liter each of NAA and BAP) as they do not grow Regeneration of Plants. Colonies are transferred directly to regenerawell as calli.

Where shoot regeneration occurs, this is manifested as a mossy type growth of many shoot primordia. On transfer to the same medium with the addition of 0.1 mg/liter GA3, these normalize, eventually to be rooted, cotted on, and moved to the greenhouse.

3. Preparation and Culture of Protoplasts from Roots of Phaseolus aureus (Mung Bean)

The use of roots as a source of protoplasts was first described in protoplasts from a number of species has been described. We describe only to root protoplasts, but also to seedling protoplasts, of a number of 960.2 However, it is only recently that their use as a source of dividing nere the protocol of Xhu et al., 28 which this group has recently applied not grain legumes and Brassica.30

Source of Material. Seeds of Phaseolus aureus cv. Pism-2 (Vigna England) for 30 min and washed 3 times with sterile tap water. Sterile radiata, ming bean) were surface sterilized with 10% v/v commercial bleach (Domestos; Lever Brothers, Kingston upon Thames, KT12BA, etri dishes seeds were germinated in the dark (27%) for 24-30 hr in 9-cm each containing 10 ml sterile tap water (50 seeds per dish).

incubated with gentle agitation on a rotatory shaker (16 hr, 25°, 60 rpm) in Preparation of Protoplasts. Root tips (1 cm long) were excised, cut solution (see the table). Plasmolysed sections from 100 root pieces were transversely into 0.5-1 mm sections, and plasmolysed for 1 hr in CPW13M 4 ml of enzyme solution (2% w/v Rhozyme, 4% w/v Meicelase, 0.03% w/v

E. C. Cocking, Nature (London) 187, 927 (1960).

²² Z.-H. Xhu, M. R. Davey, and E. C. Cocking, Z. Pflanzenphysiol, 104, 289 (1981).
²³ M. R. Davey, in "Protoplasts 1983: Lecture Proceedings (I. Potrykus, C. T. Harms, A. Hinnen, R. Hutter, P. J. King, and R. D. Shillito, eds.), p. 19. Birkhauser, Basel, 1983.

Macerozyme R10 in CPW13M). Antibiotics were included (400 mg/liter ampicillin, 10 mg/liter gentamycin, 10 mg/liter tetracycline) in the enzyme solution in some cases.

Protoplasts were released by gently squeezing. The enzyme-protoplast mixture was filtered through a nylon sieve (64 µm mesh) and the protoplasts pelleted (10 min). The pellet was resuspended in CPW21S solution (see the table) and centrifuged (5 min). Floating protoplashs were collected and washed twice with CPW13M, resuspended in culture medium, and counted.

Culture. Protoplasts were cultured at 2-2.5 \times 104/ml at 23° in the dark DK-4000, Roskilde, Denmark) each containing 2 ml of liquid medium or I ml of liquid medium over 1 ml of agar (Sigma, 0.5% w/v) solidified or diffuse light in 56 × 14 mm Nunclon dishes (A/S Nunc, Kamstrup,

A number of media have been tested. The medium given (modified B5P, see the table) gave good results. B511 medium (1 ml) (B5P containing derived colonies were collected after 15-20 days growth and replated in 3% w/v sucrose and no mannitol) was added after 7-10 days. Protoplastliquid modified White's medium2 with 0.5 mg/liter each of NAA and

4. Isolation and Culture of Protoplasts from a Cell Suspension Culture of Daucus carota (Carrot)

plasts if regeneration to plants is not necessarily required. We describe a sion cultures and are a good source of large numbers of uniform protomethod for use with cultures of carrot (in this case a nonmorphogenic Protoplasts can be isolated and cultured from a wide range of suspenline), which allows one to easily obtain 108 protoplasts in one isolation.

Source of Material. Suspension cultures of carrot3 maintained in a Murashige and Skoog¹⁴ based medium (CS; see the table) by weekly subculture (1:7 dilution) and grown on a gyratory shaker (110 rpm, 2 cm throw) at 26° in the dark.

fer are harvested by centrifugation (500 g, 5 min), resuspended in two Preparation of Protoplasts. Suspension cultures 2-3 days after transzuka" R10, 1% w/v Macerozyme R10, 0.5% w/v Driselase, in 0.4 M times the original volume of enzyme solution (2% w/v Cellulase "Onomannitol, 0.025 M CaCl, , MES, pH 5.6) and poured (10 ml/9-cm dish) into

¹¹ O. L. Gambong, R. A. Miller, and K. Ohyama, Exp. Cell Res. 59, 151 (1968).

² H. N. Wood and A. C. Braun, Proc. Natl. Acad. Sci. U.S.A. 47, 1907 (1961).

[&]quot;R. H. Smith, this volume [22].

T. Murashige and F. Skoog, Physiol. Plant. 15, 473 (1962)

petri dishes. These are sealed and incubated at 24° on a rocking table at 20-30 rpm for 4-5 hr. Preparations are checked regularly for the release

of protoplasts.

M mannitol, 0.125 M CaCl2, MES, pH 5.6), counted, and resuspended in The protoplasts are washed twice by sedimentation with osmoticum (0.29 The digest is filtered through a 50- $\mu \mathrm{m}$ stainless-steel sieve, transferred into capped centrifuge tubes, and sedimented by centrifugation for 5 min. culture medium (CP, see the table) at a population density of $2 \times 10^{5/mL}$

Culture. The protoplasts can be manipulated in various ways and then Cultures are diluted by weekly addition of an equal volume of medium cultured in CP medium at this density in liquid medium at 26° in the dark. without mannitol to reduce the osmotic pressure.

Alternatively, they can be incorporated into agar or agarose solidified medium and cultured as described for petunia and tobacco protoplasts.

Suspension Culture of a Graminaceous Species Lolium multiflorum 5. Preparation and Culture of Protoplasts from a Nonmorphogenic

which divide. There have been reports of division and colony formation of protoplasts from morphogenic suspension cultures of these species36 but general divide in culture although there have been exceptions to this Protoplasts from leaf or other whole plant tissues of grasses do not in rule.35 However, there are a number of suspension cultures of graminaceous species available, and these have been used to produce protoplasts

We describe a protocol developed in our laboratory for the isolation and culture of protoplasts from Lolium multiflorum (Italian Ryegrass) these have not yet proved to be repeatable.

suspension culture cells.

Source of Material. The cell line was originally established by P. J. Dale who has also used it for protoplast culture."

Suspension cultures are maintained by weekly serial transfer (1:7 liter 2,4-D on a gyratory shaker (110 rpm, 2 cm throw) in low light levels dilution) in CC medium38 (see the table) without mannitol and with 2 mg/

6 days after subculture. Ten milliliters of cells is sedimented by centrifugation (5 min) and resuspended in the same volume of enzyme solution Preparation of Protoplasts. Cultures are used for protoplasting 4, 5, or (4% w/v Driselase in 0.38 M mannitol, 8 mM CaCl2, MES, pH 5.6). (500 lux).

J. Potrykus, in "Advances in Protoplast Research" (L. Ferenczy and G. L. Ferkas, eds.) p. 243. Hung. Acad. Sci., Budapest, 1980.

11. Potrykus, C. T. Harms, and H. Loetz, Theor. Appl. Genet. S4, 209 (1979). M. G. K. Jones and P. J. Dale, Z. Pflanzenphysiol. 118, 267 (1982). W. Vasil and I. K. Vasil, Theor. Appl. Genet. 36, 97 (1980).

[23] PROTOPLAST CULTURE

placed on a rocking table for 1 hr at 20° before being incubated overnight (15 hr) without agitation at the same temperature. The preparation is then The solution is poured into a 9-cm petri dish and this is sealed and placed on the rocking table for an hour followed by another hour without

suspension distributed into 2 centrifuge tubes. After centrifugation to sediment the protoplasts (10 min) they are taken up in 3 ml osmoticum L1 The protoplasts are filtered through a 100-µm mesh stainless-steel sieve, an equal volume of 0.2 M CaCl₂ (MES, pH 5.8) added, and the $(0.25\,M$ mannitol, $0.1\,M$ CaCl, MES, pH 5.8) and overlayered on a 5 ml sucrose cushion (0.6 M sucrose, MES, pH 5.8).

Protoplasts collecting at the interface after centrifugation are carefully removed and washed twice with osmoticum L1, counted, and resuspended in CC medium (see the table) at a density of 2×10^6 /ml.

Culture. The protoplasts are cultured in 3.5-cm-diameter petri dishes (2 ml per dish) at 26° in the dark. Fresh CC medium with 0.2 M mannitol is added after 7 days to dilute the culture and reduce the osmotic pressure and medium without mannitol added weekly thereafter.

Calli arising from the cultures are grown on CC medium as used for the suspension cultures solidified with 0.8% w/v cleaned agar

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J

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5. The Synthesis of High-Molecular-Weight Proteins in M. the Wheat Germ Translation System G. W. W.

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[22] Establishment of Calli and Suspension Cultures¹ By ROBERTA H. SMITH

A plant callus is a wound response from an explant (the fragment of a plant or tissue used to initiate a culture) consisting of unorganized, dividing cells. Cellular proliferation can also be produced in vitro without physical injury or wounding by germinating seeds on a medium containing a plant growth regulator. Individual cells in a callus mass can vary in size, shape, pigmentation, and appearance. Most are differentiated cells with a large central vacuole and nucleus to one side as opposed to undifferentiated, meristematic cells which are isodiametric, small, lack a prominent vacuole; are cytoplasmic, and have a large central nucleus.

Callus cultures were first reported in 1939 independently by three laboratories. Gautheret¹⁶ and Nobécourt² used carrot (Daucus carota L.) root tissues with 3-indoleacetic acid (IAA) to obtain callus growth. White used procambial tissue from a tobacco hybrid (Nicotiana glauca × N. langsdorffii) to establish callus. Since then, callus cultures have been used for studies of cell division, morphogenesis, secondary product synthesis, bioassays for plant growth regulators, and more recently for selection, mutation, and genetic modification at the cellular level.

As callus cultures are subcultured (divided and placed on fresh medium, usually at 4-week intervals) many things can happen to change the original callus composition. Some of the cell types within a culture will divide faster than others and tend to overgrow and become the predominate cellular type composing a culture. Friability may change as a result. The ploidy of a culture may change with time. The cells may lose their requirement for plant growth regulators (i.e., become habituated). Secondary metabolite production can change. If plants are regenerated from successive subcultures, the incidence of variant plants (somaclonal variation)³ can increase.

Callus and suspension culture initiation involves three major considerations—selection of explant, medium, and culture conditions—each one

METHODS FOR PLANT MOLECULAR BIOLOGY

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¹ Texas Agricultural Experiment Station manuscript No. 19909

^{B.} R. J. Gautheret, C.R. Hebd. Seances Acad. Sci. 236, 118 (1939).
¹ P. Nobécourt, C.R. Seances Soc. Biol. 138, 1270 (1939).

P. R. White, Am. J. Bot. 25, 59 (1939).

T. Murashige and R. Nakano, Am. J. Bot. 54, 9763 (1967).

P. J. Larkin and W. R. Scoweroft, Theor. Appl. Genet. 60, 197 (1981).

Explant Procedure

method of surface sterilization which follows can be applied for surface Seedling tissue arising from the aseptically germinated seed is a choice source of explant tissue for carrot and tobacco callus induction. The sterilization of most explant tissue. It is more difficult to obtain clean cultures from some explants than others. The concentration of bleach and length of exposure can be varied. For a discussion of isolation of other explant sources and sterilizing agents, refer to Street,6 Yeoman,9 and Murashige. 10 Laboratory Protocol: Explant Preparation: Aseptic Germination of Seeds

Seed Germination Medium

- mination medium (see Medium Preparation section to prepare stocks), the 1. 25 \times 150 mm culture tubes or petri dishes containing 25 ml of ger-30 g sucrose, dilute to I liter, adjust pH to 5.7, add 8 g of agar, and heat composition of which is as follows: 10 ml each of the inorganic salt stocks, until agar dissolves; pour 25 ml each into tubes or petri dishes.
 - 2. Rinse water (200 ml) in a 500-ml Erlenmeyer flask, capped with aluminum foil
- 3. Petri dishes in a cantister.
- 6 H. E. Street, in "Plant Tissue and Cell Culture" (H. E. Street, ed.), p. 1. Univ. of California Press, Berkeley, 1973.
 - T. Murashige, TCA Rep. 12(2), 41 (1978).
 - Murashige, HortScience 12(2), 3 (1977).
- M. M. Yeoman, in "Flant Tissue and Cell Culture" (H. B. Street, ed.), p. 31. Univ. of California Press, Berkeley, 1973.
 - 4 T Mirrathine. Annu. Rev. Plant Physiol. 25, 135 (1974).

[22] CALLI AND SUSPENSION CULTURES

4. Autoclave seed germination medium for 15 min and rinse water and petri dishes for 20 min at 121°.

Sterilization of Seeds

Carrot

- 1. Wrap the seeds in several layers of ca. 10×10 -cm squares of cheesecloth.
 - 2. Place the wrapped seeds in a 25×150 -mm culture tube, add 70% ethanol, shake or swirl for ~ 2 min, and then decant the alcohol.
- 3. Disinfect the wrapped seeds in a 20% (v/v) solution of commercial bleach or in a solution composed of 1.05% sodium hypochlorite and 2 drops of Tween-20 per 100 ml total volume. Pour the disinfectant into the cap of the test tube and then into the culture tube, completely covering the cheesecloth-wrapped seeds. Cap the tube.
 - 4. Swirl the tube intermittently on a vortex mixer for 20 min. Using aseptic procedures, place the materials in a laminar air flow hood, pour off the disinfectant, and rinse the seeds 3 times with sterile water.
 - Note: Sometimes carrot seeds are very contaminated: therefore, it is prudent to plant only 2-3 seeds per dish or tube. Seed contamination will be apparent on the seed germination medium after about 5 days.

Tobacco

- 1. Put the seeds in a 15-ml screw-cap, conical centrifuge tube and add 0.75% sodium hypochlorite (containing 2 drops Tween-20/100 ml).
- 2. Swirl the tube intermittently on a vortex mixer for 20 min.
 - 3. Allow the seeds to settle or centrifuge at low speed.
- 4. Pour off the disinfecting solution and rinse 3 times with sterile

Seed Germination

- 1. Using long-handled forceps (Pott Smith dressing forceps), plant 1-3 seeds per culture tube on the surface of the agar. To sterifize forceps, immerse in 70% alcohol 10 min. Then place tips in 95% alcohol covering and pass forceps tips through a flame. Allow alcohol to burn off and forceps to cool prior to touching plant tissue. (Caution: hold forceps close ca. one-half the length of the forceps. Prior to using, remove from alcohol to horizontal so flame does not burn hand.) Note: Many Seginners will kill the plant tissue by not allowing the instruments to cool
- 2. Place cultures in 16:8 hr (light: dark) photoperiod at light intensity of 50-100 μ Em⁻² sec⁻¹ and temperature 26 ± 3°.

Fro. 1. Seedlings of carrot (left) and tobacco (right) aseptically germinated.

3. In 1-3 weeks seedlings from aseptically germinated seeds will be ready for callus induction; all parts of the seedlings can be used as an explant for callus induction (Fig. 1).

Nutrient Medium

Components of a nutrient medium for callus growth generally include organic supplements (sometimes optional). Inorganic salts, vitamins, and inorganic salts, a carbohydrate, vitamins, plant growth regulators, and plant growth regulator stock solutions are prepared in concentrated form, and usually several inorganic salts and vitamins can be combined to minimize the number of stocks. Plant cell culture medium is a rich substrate which will support growth of bacteria and fungi. Generally, addition of fungicides and bactericides to medium can result in complications and eration. Always use glass-distilled or demineralized water to prepare stocks and media. Detailed discussions of common media are proved ciency of medium preparation. Storage of saft stocks is best under refrigphytotoxicity to the explant. 11 Stock solutions enhance accuracy and effielsewhere. 12-16

[22] CALLI AND SUSPENSION CULTURES

Laboratory Protocol: Medium Preparation

Stock Solutions

1. Inorganic salts of Murashige and Skoog, 15 100 times final medium concentration (grams/liter stock):

CuSO, 5H2O, 0.0025 ZaSO, · 7H2O, 0.860 MnSO₄ · H₂O, 1.690 MgSO4 - 7H2O, 37 NH,NO3, 165 KNO, 190 Nitrates Sulfates Halides

CaCl₂ · 2H₂O, 44 KI. 0.083

CoCl. 6H20, 0.0025 PO4, BO3, MoO4 H,BO, 0.620 KH₂PO₄, 17

NaMoO4 · 2H2O, 0.025 NaFe EDTA

FeSO₄ · 7H₂O, 2.784 Na₂EDTA, 3.724

Store in amber bottle

phenoxyacetic acid (2,4D) and IAA stocks. Weigh out 10 mg auxin into a 2. Plant growth regulators (store in refrigerator): (1) 2,4-dichloro-200 mi beaker, add several drops of 1 N NaOH or KOH until crystals are dissolved (not more than 0.3 ml), rapidly add 90 ml of double distilled H₂O, bring up to 100 ml volume in a volumetric flask. Make IAA stock fresh weekly. (2) Kinetin stock: same as for auxins except use 1 N HCI and a few drops of H₂O with gentle heating to dissolve crystals.

3. Vitamins (store in freezer in 10 ml aliquots) (mg/100 ml stock): Pyridoxine-HCl, 5 Nicotinic acid, 5 Thiamin-HCl, 4

¹¹ K. C. Thurston, S. J. Spencer, and U. Ardini, Am. J. Bot. 66, 825 (1979).

¹¹ O. L. Gamborg, R. A. Miller, and K. Ojima, Exp. Cell Rev. 59, 151 (1969).
¹¹ O. L. Gambore T. Mirrachine, T. A. Thorne and I. K. Vacil In Viton 19, 474 (1976).

¹⁴ L. C. Huang and T. Murashipe, Tissue Cult. Assoc. Man. 3, 539 (1977)

¹⁶ T. Murashige, in "Tissue Culture Methods and Applications". (P. F. Kruse and M. K. 13 T. Murashige and F. Skoog, Physiol. Plant. 15, 473 (1962).

Patterson, Jr., eds.), p. 698. Academic Press, New York, 1973.

Medium, I Liter

1. In a 2-liter Erlenmeyer flask add 500 ml of deionized, distilled water and mix in the following:

Carrol 17-19

- 10 ml each of the five inorganic salt stocks
- 30 g sucrose
- 10 ml vitamin stock
- 100 mg myo-inositol
- 1 ml 2,4-D stock, (0.5 µM)

Tobacco 6.15.18-20

- 10 ml each of the inorganic salt stocks
- 30 g sucrose
- 10 ml vitamins
- 100 mg myo-inositol
- 2 ml kinetin stock (0.9 μM)
- 20 ml IAA stock (11.4 μM)
- (Leaf or stem explants of Nicotiana spp. have been used to initiate callus on MS medium with 4.5 μM 2,4-D and 2 g/l casein hydroly-
- 2. Final pH adjusted to 5.7 with 1 N KOH, NaOH, or HCl. Add 8 g of

Heat and stir while melting the agar so that the agar does not burn on the bottom of the flask. Distribute 25 ml/culture tube. An alternate procedure to melt the agar is to autoclave the flask 3-7 min at 121°.14 After melting and autoclaving (15 min at 121°C), distribute 25 ml per sterile, plastic petri dish in a laminar air flow hood.

Care should be taken in autoclaving media. Sugars will undergo caramelization if autoclaved too long, and will also react with amino compounds (Maillard Reaction).21 These then form brown, high-molecularweight materials which can inhibit growth. A brown color to the medium

- "D. F. Wetherell, in "Propagation of Higher Plants through Tissue Culture" (K. W. Hughes, R. Henke, and M. Constantin, eds.), p. 102. Tech. Inf. Cent., USDB, 1978.
- 19 D. A. Evana, W. R. Sharp, and C. E. Flick, in "Horricultural Reviews" (J. Janik, ed.), p.
- 9 O. L. Gamborg and L. R. Wetter, in "Plant Tissue Culture Methods" (O. L. Gamborg and 214. AVI Publ. Co., Westport, Connecticut, 1980.
- w J. P. Helgeson, to "Nicotiana: Procedures for Experimental Use" (R. D. Durbin, ed.), L. R. Wetter, eds.), p. 95. Natl. Res. Counc. Can., 1975.
 - U.S. Dept. Agric., Tech. Bull. No. 1586, p. 52. USDA, Washington, D.C., 1979.
- 11 H. G. Peer, in "Effects of Sterilization on Components in Nutrient Media" (J. van Bragt, D. A. A. Mossel, P. L. M. Pierik, and H. Vetústra, eds.); Misc. Pap. No. 9, p. 105.



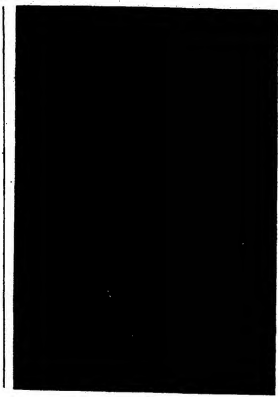


Fig. 2. Carrot explants ready to be placed in culture.

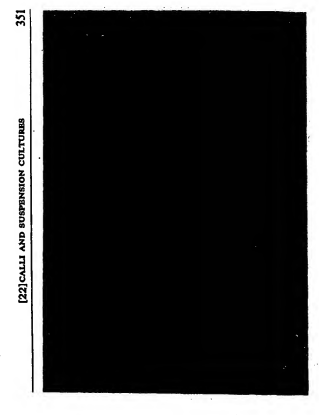
is apparent if media is autoclaved too long or left in the autoclave to cool down. Media autoclaving should be accurately timed, and removed from the autoclave promptly when the temperature is just below 100° and pressure is zero.

Media can be stored 1-2 weeks prior to use, preferably under refrigeration in the dark. Media with IAA cannot be stored as IAA breaks down.

Laboratory Protocol: Culture of Primary Explants for Callus Induction

- 1. Carrot explants approximately 1 cm long from seedling roots and petioles are excised and placed on the medium (Fig. 2)
 - 2. Tobacco explants from cross-sections of leaf and stem tissue are excised and placed on the medium.
- 3. All cultures are incubated in the dark at 27°.
- 4. After 4-6 weeks callus can be separated from the original explant and subcultured onto fresh medium for further growth (Fig. 3). The stock callus cultures should be subdivided and cultured at 4- to 6-week intervals

Sometimes after 3-5 days in culture, bacterial and/or fungal growth will he annarent. Contamination that results from the explant (spores on



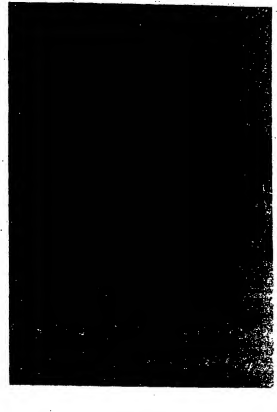


Fig. 4. Carrot (top) and tobacco (bottom) stock callus cultures.

FIG. 3. Carrot (top) and tobacco (bottom) callus beginning to form from the explants after about a week in culture. tissue, rinse water, petri dish, instruments) will arise on the medium adjacent to the explant. Contamination scattered on the agar surface usually results from improperly sterilized media, contaminated air flow hood,

For details on media for callus induction for various other dicots and monocots (see Evans et al. 18). or poor sterile technique.

Suspension Culture

Suspension cultures (cell cultures dispersed in moving liquid medium)⁶ are established from friable callus cultures. Friable callus can be visually selected and subcultured from compact, hard cultures or can be achieved

Fig. 5. Carrot suspension culture after about 10 days growth.

by manipulating medium components including the type or concentration of auxin, 22 cytokinin, or casein hydrolysate. 8 Environmental culture conditions may also influence friability.

medium except lacking agar. Baffled, long-necked Bhrlenmeyer flasks are Suspension cultures of carrot (Fig. 5) can be initiated by subculturing commonly used and placed on an orbital or gyratory shaker at 50-120 rpm. Tobacco suspension cultures can be cultured on the same callus induction basal medium, substituting 1 μM 2,4-D for IAA and deleting the carrot callus onto liquid medium of the same composition as callus

Suspension cultures are initiated by an inoculum which establishes a cell density of 0.5-2.5 \times 10⁵ cells per ml. ²³ If too low a density inoculum is used, the culture may not grow. Generally, 2-3 g of friable callus is after 2 to 3 weeks. Suspension cultures require more frequent subcultures at 1- to 3-week intervals, depending on the amount of callus used to inoculated onto 100 mi of liquid medium. Small clumps of cells will result

initiate the culture and the vigor of the culture. A pipette with a wide tip can be used to inoculate new cultures or the old suspension culture can be swirled and a specified volume rapidly poured into a sterile, graduate dium. According to Helgeson, 20 if cell concentrations exceed 40 mg fresh weight per milliliter of medium, transfers with no appreciable lag before the logarithmic phase can be established. A fine dispersion of single cells and small cell chusters can be established and maintained by screening the clumps. Agar medium cultures can be initiated from suspension cultures by pipetting 1-3 ml of suspension culture onto the surface of an agarevlinder. The contents of the cylinder is then poured into the new mesuspension culture through a nylon or wire mesh to exclude larger cell solidified petri dish.

Growth Curve

For either a suspension or callus culture it is important to establish a growth curve. Growth curves can be established by inoculating cultures with a specified volume or weight of cells and then taking daily interval measurements. Fresh weight, dry weight, and mitotic index 2.19,22 can be used to follow culture growth. Suspension-culture-packed cell volume (volume of cells per unit volume of culture) measurement is also a rapid method to follow growth.2 A specified aliquot of suspension culture is centrifuged in a 15-ml graduated conical centrifuge tube at 2000 g for 5 min. A measurement in milliliters of the packed cell volume per milliliter culture can be followed. A growth curve with cell number, weight, etc., on the y axis and time intervals on the x axis will have 5 segments: the lag, exponential, linear, progressive decelerating, and stationary.23 Generally, a culture should be subcultured prior to reaching stationary phase.

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